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present
NEWS 4 AUG 05 New pricing for EUROPATFULL and PCTFULL effective
August 1, 2003
NEWS 5 AUG 13 Field Availability (/FA) field enhanced in BEILSTEIN
NEWS 6 AUG 18 Data available for download as a PDF in RDISCLOSURE
NEWS 7 AUG 18 Simultaneous left and right truncation added to PASCAL
NEWS 8 AUG 18 FROSTI and KOSMET enhanced with Simultaneous Left and Right
Truncation
NEWS 9 AUG 18 Simultaneous left and right truncation added to ANABSTR
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AND CURRENT DISCOVER FILE IS DATED 23 SEPTEMBER 2003
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FILE 'HOME' ENTERED AT 14:37:11 ON 04 NOV 2003

=> file reg

COST IN U.S. DOLLARS

SINCE FILE

TOTAL

ENTRY

SESSION

FULL ESTIMATED COST

0.21

0.21

Patel

<11/4/2003>

FILE 'REGISTRY' ENTERED AT 14:37:20 ON 04 NOV 2003
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STRUCTURE FILE UPDATES: 3 NOV 2003 HIGHEST RN 612478-18-9
DICTIONARY FILE UPDATES: 3 NOV 2003 HIGHEST RN 612478-18-9

TSCA INFORMATION NOW CURRENT THROUGH JULY 14, 2003

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Crossover limits have been increased. See HELP CROSSOVER for details.

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=>
Uploading 10046526.3

L1 STRUCTURE UPLOADED

=> s l1
SAMPLE SEARCH INITIATED 14:37:48 FILE 'REGISTRY'
SAMPLE SCREEN SEARCH COMPLETED - 20 TO ITERATE

100.0% PROCESSED 20 ITERATIONS 0 ANSWERS
SEARCH TIME: 00.00.01

FULL FILE PROJECTIONS: ONLINE **COMPLETE**
BATCH **COMPLETE**
PROJECTED ITERATIONS: 132 TO 668
PROJECTED ANSWERS: 0 TO 0

L2 0 SEA SSS SAM L1

=> s l1 sss full
FULL SEARCH INITIATED 14:37:54 FILE 'REGISTRY'
FULL SCREEN SEARCH COMPLETED - 326 TO ITERATE

100.0% PROCESSED 326 ITERATIONS 1 ANSWERS
SEARCH TIME: 00.00.01

L3 1 SEA SSS FUL L1

=> file marpat
COST IN U.S. DOLLARS SINCE FILE TOTAL
ENTRY SESSION
FULL ESTIMATED COST 148.15 148.36

FILE 'MARPAT' ENTERED AT 14:38:09 ON 04 NOV 2003

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FILE CONTENT: 1988-PRESENT (VOL 104 ISS 15-VOL 139 ISS18) (20031101ED)

MOST RECENT CITATIONS FOR PATENTS FROM FIVE MAJOR ISSUING AGENCIES
(COVERAGE TO THESE DATES IS NOT COMPLETE):

US 6627682 30 SEP 2003
DE 20300703 25 SEP 2003
EP 1350541 08 OCT 2003
JP 2003286251 10 OCT 2003
WO 2003082838 09 OCT 2003

Structure search limits have been raised. See HELP SLIMIT for the new,
higher limits.

=> s l1 sss full

FULL SEARCH INITIATED 14:38:15 FILE 'MARPAT'
FULL SCREEN SEARCH COMPLETED - 11133 TO ITERATE

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95.3% PROCESSED	10611 ITERATIONS (1 INCOMPLETE)	1 ANSWERS
99.4% PROCESSED	11067 ITERATIONS (5 INCOMPLETE)	5 ANSWERS
100.0% PROCESSED	11133 ITERATIONS (5 INCOMPLETE)	5 ANSWERS

SEARCH TIME: 00.01.31

L4 5 SEA SSS FUL L1

=> d his

(FILE 'HOME' ENTERED AT 14:37:11 ON 04 NOV 2003)

FILE 'REGISTRY' ENTERED AT 14:37:20 ON 04 NOV 2003

L1 STRUCTURE UPLOADED
L2 0 S L1
L3 1 S L1 SSS FULL

FILE 'MARPAT' ENTERED AT 14:38:09 ON 04 NOV 2003

L4 5 S L1 SSS FULL

=> file caplus

COST IN U.S. DOLLARS	SINCE FILE ENTRY	TOTAL SESSION
FULL ESTIMATED COST	105.35	253.71

FILE 'CAPLUS' ENTERED AT 14:40:03 ON 04 NOV 2003

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FILE COVERS 1907 - 4 Nov 2003 VOL 139 ISS 19
FILE LAST UPDATED: 3 Nov 2003 (20031103/ED)

This file contains CAS Registry Numbers for easy and accurate substance identification.

=> d 15 fbib hitstr abs total

L5 NOT FOUND

The L-number entered has not been defined in this session, or it has been deleted. To see the L-numbers currently defined in this session, enter DISPLAY HISTORY at an arrow prompt (=>).

=> s 13

L5 2 L3

=> s 14

L6 5 L4

=> s 15 and 16

L7 0 L5 AND L6

=> d 15 fbib hitstr abs total

L5 ANSWER 1 OF 2 CAPLUS COPYRIGHT 2003 ACS on STN

AN 2003:551181 CAPLUS

DN 139:117339

TI Preparation of substituted arylamine derivatives as antitumor agents

IN Elbaum, Daniel; Askew, Benny; Booker, Shon; Germain, Julie; Habgood, Gregory; Handley, Michael; Kim, Tae-Seong; Li, Aiwen; Nishimura, Nobuko; Patel, Vinod F.; Yuan, Chester Chenguang; Kim, Joseph L.

PA Amgen Inc., USA

SO U.S. Pat. Appl. Publ., 106 pp., Cont.-in-part of U.S. Ser. No. 46,526.

CODEN: USXXCO

DT Patent

LA English

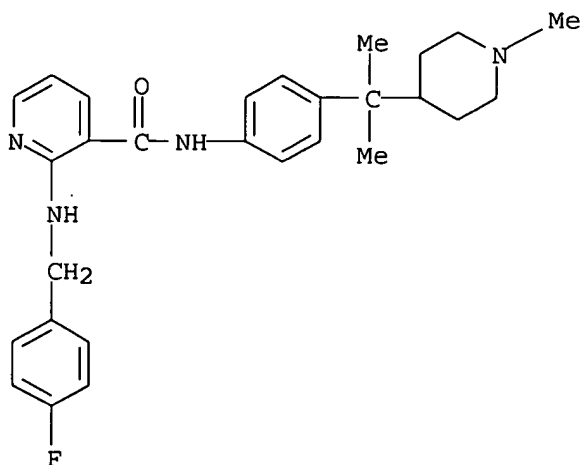
FAN.CNT 2

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
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				US 2001-323686PP	20010919
				US 2002-46526	A220020110
	US 2002147198	A1	20021010	US 2002-46526	20020110
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				US 2001-323686PP	20010919

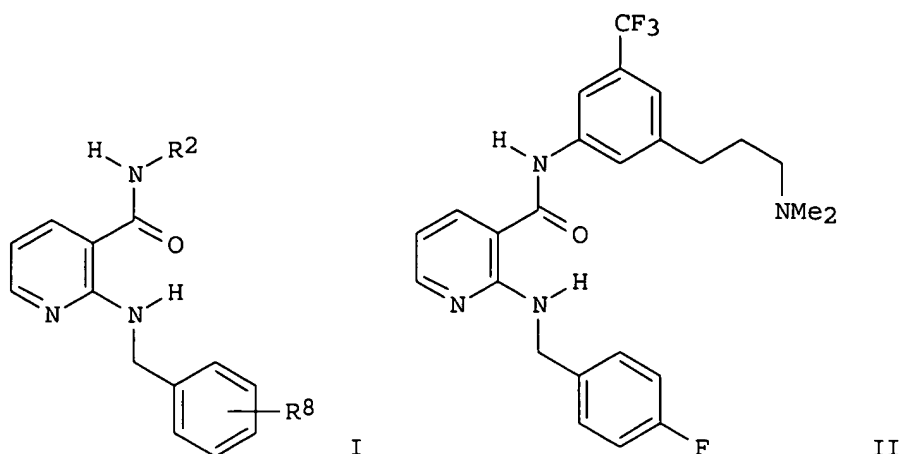
PATENT FAMILY INFORMATION:

FAN 2002:539663

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
PI	WO 2002055501	A2	20020718	WO 2002-US742	20020111
	WO 2002055501	A3	20021219		
	W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, OM, PH, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TN, TR, TT, TZ, UA, UG, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG				
				US 2001-261360PP	20010112
				US 2001-323686PP	20010919
				US 2002-46526 A	20020110
	US 2002147198	A1	20021010	US 2002-46526	20020110
				US 2001-261360PP	20010112
				US 2001-323686PP	20010919
OS	MARPAT 139:117339				
IT	442846-35-7P				
	RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses) (target compd.; prepn. of substituted aminopyridines as antitumor agents)				
RN	442846-35-7 CAPLUS				
CN	3-Pyridinecarboxamide, 2-[[[(4-fluorophenyl)methyl]amino]-N-[4-[1-methyl-1-(1-methyl-4-piperidinyl)ethyl]phenyl]- (9CI) (CA INDEX NAME)				



GI



AB The title compds. I [R2 = (un)substituted Ph, 9-10 membered bicyclic and 11-14 membered tricyclic (un)satd. heterocyclyl; R8 = halo, NH2, NO2, etc.], and their pharmaceutically acceptable derivs., are prepd. and disclosed as agents effective for prophylaxis and treatment of diseases, such as angiogenesis mediated diseases. E.g., a multi-step synthesis of II, starting from 1-dimethylamino-2-propyne and 3-bromo-5-trifluoromethylaniline, was given. Selected compds. of the invention, e.g., II, inhibited VEGF-stimulated cell proliferation at a level below 50 nM. The invention encompasses novel compds., analogs, prodrugs and pharmaceutically acceptable derivs. thereof, pharmaceutical compns. and methods for prophylaxis and treatment of diseases and other maladies or conditions involving, cancer and the like.

L5 ANSWER 2 OF 2 CAPLUS COPYRIGHT 2003 ACS on STN

AN 2002:539663 CAPLUS

DN 137:109210

TI Preparation of substituted arylamine derivatives and methods of use as antitumor agents

IN Chen, Guoqing; Booker, Shon; Cai, Guolin; Croghan, Michael; Dipietro, Lucian; Dominguez, Celia; Elbaum, Daniel; Germain, Julie; Huang, Qi; Kim, Joseph L.; Kim, Tae-Seong; Patel, Vinod F.; Smith, Leon M.; Tasker, Andrew; Xi, Ning; Xu, Shimin; Yuan, Chester Chenguang

PA Amgen Inc., USA

SO PCT Int. Appl., 253 pp.

CODEN: PIXXD2

DT Patent

LA English

FAN.CNT 2

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
PI	WO 2002055501	A2	20020718	WO 2002-US742	20020111
	WO 2002055501	A3	20021219		
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RW:	GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AT, BE, CH,				

CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR,
BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG

US 2002147198 A1 20021010

US 2001-261360PP 20010112
US 2001-323686PP 20010919
US 2002-46526 A 20020110
US 2002-46526 20020110
US 2001-261360PP 20010112
US 2001-323686PP 20010919

PATENT FAMILY INFORMATION:

FAN 2003:551181

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
PI	US 2003134836	A1	20030717	US 2002-197960	20020717
				US 2001-261360PP	20010112
				US 2001-323686PP	20010919
				US 2002-46526 A2	20020110
	US 2002147198	A1	20021010	US 2002-46526	20020110
				US 2001-261360PP	20010112
				US 2001-323686PP	20010919

OS MARPAT 137:109210

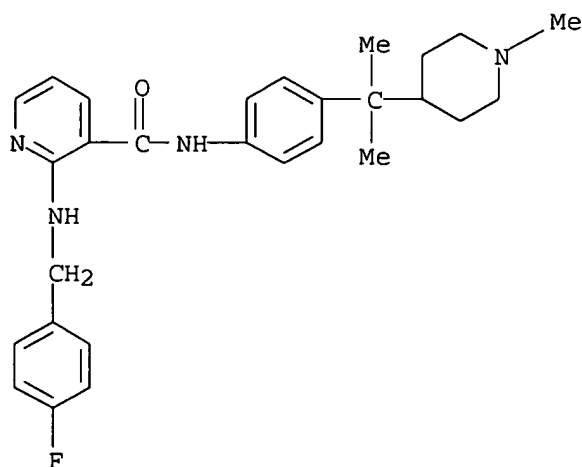
IT 442846-35-7P

RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU
(Therapeutic use); BIOL (Biological study); PREP (Preparation); USES
(Uses)

(target compd.; prepn. of substituted aminopyridines as antitumor
agents)

RN 442846-35-7 CAPLUS

CN 3-Pyridinecarboxamide, 2-[[[4-fluorophenyl)methyl]amino]-N-[4-[1-methyl-1-(1-methyl-4-piperidinyl)ethyl]phenyl]- (9CI) (CA INDEX NAME)



GI

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

AB Title compds. I [B1 and B2 independently equal C or N, wherein B1B2 form
part of 5-6 membered heteroaryl ring A; R1 = one or more substituents

selected from H, halo, oxo, (un)substituted cycloalkyl, phenylalkyl, etc.; R2 = (un)substituted cycloalkyl, cycloalkenyl, 6-10 membered aryl or 5-6 membered heterocyclyl, etc.; R3 = (un)substituted aryl; R4 = H, alkyl, (un)substituted Ph or aralkyl; X1 = bond, alkylene, alkenylene and alkynylene, where one of the CH₂ groups may be substituted with O or NH, wherein X1 is optionally substituted with OH; X2 = (un)substituted N contg. linker, e.g., -NHCH₂-, and there pharmaceutically acceptable derivs., are prepd. and disclosed as agents effective for prophylaxis and treatment of diseases, such as angiogenesis mediated diseases. Thus, II was prepd. via arylation of 1-dimethylamino-2-propyne with 3-bromo-5-trifluoromethylaniline, hydrogenation, amidation with 2-chloropyridine-3-carbonyl chloride and chloro-substitution with 4-fluorobenzylamine. Selected compds. of the invention, e.g., II, inhibited VEGF-stimulated cell proliferation at a level below 50 nM. The invention encompasses novel compds., analogs, prodrugs and pharmaceutically acceptable derivs. thereof, pharmaceutical compns. and methods for prophylaxis and treatment of diseases and other maladies or conditions involving, cancer and the like.

=> d his

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FILE 'REGISTRY' ENTERED AT 14:37:20 ON 04 NOV 2003

L1 STRUCTURE UPLOADED

L2 0 S L1

L3 1 S L1 SSS FULL

FILE 'MARPAT' ENTERED AT 14:38:09 ON 04 NOV 2003

L4 5 S L1 SSS FULL

FILE 'CAPLUS' ENTERED AT 14:40:03 ON 04 NOV 2003

L5 2 S L3

L6 5 S L4

L7 0 S L5 AND L6

=> d 14 fbib hitstr abs total

YOU HAVE REQUESTED DATA FROM FILE 'MARPAT' - CONTINUE? (Y)/N:n

=> d 16 fbib hitstr abs total

L6 ANSWER 1 OF 5 CAPLUS COPYRIGHT 2003 ACS on STN

AN 2002:290791 CAPLUS

DN 136:309922

TI Preparation of benzoxazolyl piperidines and analogs as rotamase enzyme inhibitors

IN Kemp, Mark Ian; Palmer, Michael John; Sanner, Mark Allen; Wythes, Martin James

PA Pfizer Inc, USA

SO U.S., 43 pp.

CODEN: USXXAM

DT Patent

LA English

FAN.CNT 2

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
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PI	US 6372736	B1	20020416	US 1999-358107	19990721
	US 6562964	B1	20030513	US 2002-56901	20020123
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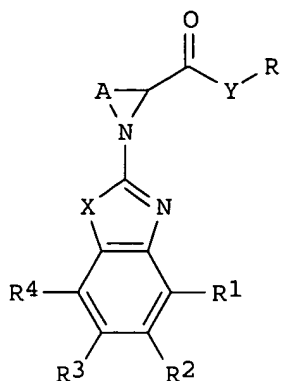
PATENT FAMILY INFORMATION:

FAN 2000:84802

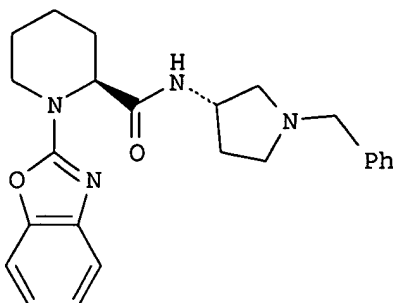
PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 2000005232	A1	20000203	WO 1999-IB1211	19990628
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RW: GH, GM, KE, LS, MW, SD, SL, SZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG				
CA 2338214	AA	20000203	GB 1998-15880	A 19980721
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AU 9942858	A1	20000214	WO 1999-IB1211 W	19990628
			AU 1999-42858	19990628
			GB 1998-15880	A 19980721
BR 9912330	A	20010417	WO 1999-IB1211 W	19990628
			BR 1999-12330	19990628
			GB 1998-15880	A 19980721
EP 1100797	A1	20010523	WO 1999-IB1211 W	19990628
EP 1100797	B1	20030226	EP 1999-963123	19990628
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			GB 1998-15880	A 19980721
EE 200100044	A	20020617	WO 1999-IB1211 W	19990628
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			GB 1998-15880	A 19980721
JP 2002521382	T2	20020716	WO 1999-IB1211 W	19990628
			JP 2000-561188	19990628
			GB 1998-15880	A 19980721
NZ 508838	A	20021220	WO 1999-IB1211 W	19990628
			NZ 1999-508838	19990628
			GB 1998-15880	A 19980721
AT 233261	E	20030315	WO 1999-IB1211 W	19990628
			AT 1999-963123	19990628
			GB 1998-15880	A 19980721
ES 2191484	T3	20030901	WO 1999-IB1211 W	19990628
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			GB 1998-15880	A 19980721
NO 2001000322	A	20010315	NO 2001-322	20010119
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HR 2001000052	A1	20011231	WO 1999-IB1211 W	19990628
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BG 105254	A	20011031	WO 1999-IB1211 W	19990628
			BG 2001-105254	20010214
			GB 1998-15880	A 19980721
			WO 1999-IB1211 W	19990628

OS MARPAT 136:309922

GI



I



II

AB Title compds. [I; A = (un)substituted unbranched C3-C5 alkylene; X and Y = independently O, S, NH, or N-alkyl; R = (un)substituted, C-linked, 4- to 6-membered, non-arom., heterocyclic ring contg. 1 N; R1-R4 = independently H, halo, (cyclo)alkyl, haloalkyl, (cyclo)alkoxy, CONR5R6, cycloalkylalkylene, cycloalkylalkoxy, or CO2R7; R5 and R6 = independently H, alkyl, or taken together = unbranched alkylene; R7 = alkyl] were prepd. as rotamase enzyme inhibitors, particularly FKBP-12 and FKBP-52 inhibitors. Thus, (2S)-1-(1,3-benzoxazol-2-yl)-2-piperidinecarboxylic acid (prepn. given) was amidated with (3S)-1-benzylpyrrolidine-3-ylamine in the presence of 1-hydroxybenzotriazole hydrate and 1-(3-dimethylaminopropyl)-3-ethylcarbodiimide.HCl in CH2Cl2 to yield II. Twenty-one compds. of the invention demonstrated inhibitory activity against human recombinant FKBP-12 in a coupled colorimetric PPIase in vitro assay with IC50 values below 1200 nM, and II inhibited the rotamase enzyme FKBP-52 in a similar assay with IC50 = 2790 nM. As neurotrophic agents, the invention compds. promote neuronal regeneration and outgrowth and are useful for the treatment of neurodegenerative diseases or other disorders involving nerve damage.

RE.CNT 20 THERE ARE 20 CITED REFERENCES AVAILABLE FOR THIS RECORD
ALL CITATIONS AVAILABLE IN THE RE FORMAT

L6 ANSWER 2 OF 5 CAPLUS COPYRIGHT 2003 ACS on STN

AN 2000:277965 CAPLUS

DN 132:308336

TI Novel imidazoles with anti-inflammatory activity and their preparation and use

IN Almansa, Carmen; Gonzalez, Concepcion; Torres, M<<fml Carmen

PA J. Uriach & Cia, S.A., Spain

SO PCT Int. Appl., 63 pp.

CODEN: PIXXD2

DT Patent

LA Spanish

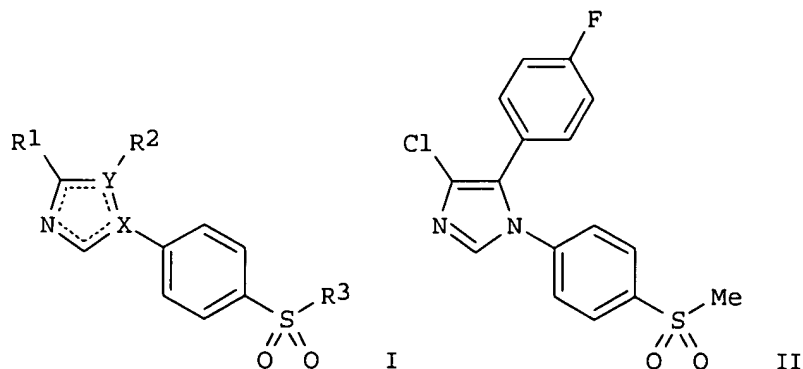
FAN.CNT 1

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 2000023426	A1	20000427	WO 1999-ES327	19991015
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MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM
 RW: GH, GM, KE, LS, MW, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG

CA 2346847	AA	20000427	ES 1998-2222 A 19981016 CA 1999-2346847 19991015 ES 1998-2222 A 19981016 WO 1999-ES327 W 19991015
AU 9962048	A1	20000508	AU 1999-62048 19991015 ES 1998-2222 A 19981016 WO 1999-ES327 W 19991015
EP 1122243	A1	20010808	EP 1999-949025 19991015
R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO			ES 1998-2222 A 19981016 WO 1999-ES327 W 19991015
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JP 2002527508	T2	20020827	JP 2000-577154 19991015 ES 1998-2222 A 19981016 WO 1999-ES327 W 19991015
NO 2001001790	A	20010614	NO 2001-1790 20010409 ES 1998-2222 A 19981016 WO 1999-ES327 W 19991015

OS MARPAT 132:308336
 GI



AB Title compds. I are disclosed [wherein one of X or Y = N; other = C; R1 = H, Me, halo, cyano, NO₂, CHO, COCH₃ or COOR₄; R2 = (un)substituted aryl or heteroaryl; R3 = C1-8 alkyl, C1-8 haloalkyl, or NR₄R₆; R4 = H, C1-8 alkyl, or (un)substituted aryl(alkyl); R6 = H, C1-8 alkyl, arylalkyl, COR₈, or CO₂R₈; R8 = C1-8 alkyl or C1-8 haloalkyl; aryl = Ph or naphthyl; heteroaryl = pyridyl, pyrazinyl, pyrimidinyl, or pyridazinyl, optionally fused to a benzene ring]. The compds. are useful as selective inhibitors of cyclooxygenase-2 (COX-2), and particularly as antiinflammatories. Claimed uses include treatment of inflammation, pain, fever,

prostanoid-induced smooth muscle contraction, dysmenorrhea, premature labor, asthma, bronchitis, cancer (esp. gastrointestinal or colon), cerebral infarct, epilepsy, or neurodegenerative diseases such as Alzheimer's disease or dementia. For instance, 4-(MeSO₂)C₆H₄NH₂ was condensed with 4-FC₆H₄CHO under Dean-Stark conditions, and the resulting imine was cyclized with tosylmethyl isocyanide (75%), followed by imidazole ring chlorination with N-chlorosuccinimide (80%), to give the invention compd. II. This compd. gave 89% inhibition of COX-2 at 1 .mu.M in vitro, but only 37.8% inhibition of COX-1 at 10 .mu.M.

RE.CNT 5 THERE ARE 5 CITED REFERENCES AVAILABLE FOR THIS RECORD
ALL CITATIONS AVAILABLE IN THE RE FORMAT

L6 ANSWER 3 OF 5 CAPLUS COPYRIGHT 2003 ACS on STN
AN 2000:84802 CAPLUS
DN 132:137377
TI Preparation of benzoxazolyl piperidines and analogs as rotamase enzyme inhibitors
IN Kemp, Mark Ian; Palmer, Michael John; Sanner, Mark Allen; Wythes, Martin James
PA Pfizer Limited, UK; Pfizer Inc.
SO PCT Int. Appl., 131 pp.
CODEN: PIXXD2
DT Patent
LA English
FAN.CNT 2

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
PI	WO 2000005232	A1	20000203	WO 1999-IB1211	19990628
	W:	AE, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ, DE, DK, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, UA, UG, US, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM			
	RW:	GH, GM, KE, LS, MW, SD, SL, SZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG			
	CA 2338214	AA	20000203	GB 1998-15880 A 19980721 CA 1999-2338214 19990628 GB 1998-15880 A 19980721 WO 1999-IB1211 W 19990628	
	AU 9942858	A1	20000214	AU 1999-42858 19990628 GB 1998-15880 A 19980721 WO 1999-IB1211 W 19990628	
	BR 9912330	A	20010417	BR 1999-12330 19990628 GB 1998-15880 A 19980721 WO 1999-IB1211 W 19990628	
	EP 1100797	A1	20010523	EP 1999-963123 19990628	
	EP 1100797	B1	20030226		
	R:	AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, PT, IE, SI, LT, LV, FI, RO			
	EE 200100044	A	20020617	GB 1998-15880 A 19980721 WO 1999-IB1211 W 19990628 EE 2001-44 19990628 GB 1998-15880 A 19980721 WO 1999-IB1211 W 19990628	
	JP 2002521382	T2	20020716	JP 2000-561188 19990628 GB 1998-15880 A 19980721	

NZ 508838	A	20021220	WO 1999-IB1211 W 19990628
			NZ 1999-508838 19990628
			GB 1998-15880 A 19980721
AT 233261	E	20030315	WO 1999-IB1211 W 19990628
			AT 1999-963123 19990628
			GB 1998-15880 A 19980721
ES 2191484	T3	20030901	WO 1999-IB1211 W 19990628
			ES 1999-963123 19990628
NO 2001000322	A	20010315	GB 1998-15880 A 19980721
			NO 2001-322 20010119
			GB 1998-15880 A 19980721
HR 2001000052	A1	20011231	WO 1999-IB1211 W 19990628
			HR 2001-52 20010119
			GB 1998-15880 A 19980721
BG 105254	A	20011031	WO 1999-IB1211 W 19990628
			BG 2001-105254 20010214
			GB 1998-15880 A 19980721
			WO 1999-IB1211 W 19990628

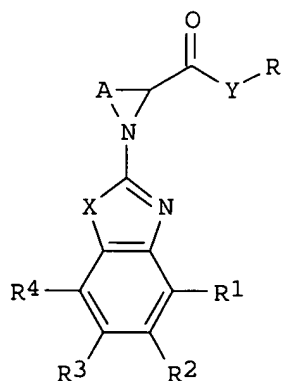
PATENT FAMILY INFORMATION:

FAN 2002:290791

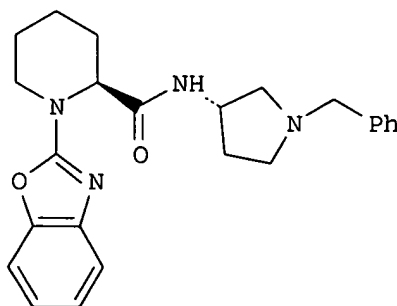
	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
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PI	US 6372736	B1	20020416	US 1999-358107	19990721
	US 6562964	B1	20030513	US 2002-56901	20020123
				GB 1998-15880 A	19980721
				US 1999-358107 A3	19990721

OS MARPAT 132:137377

GI



I



II

AB Title compds. (I) [wherein A = (un)substituted unbranched C3-C5 alkylene; X and Y = independently O, S, NH, or N-alkyl; R = (un)substituted, C-linked, 4- to 6-membered, non-arom., heterocyclic ring contg. 1 N; R¹-R⁴ = independently H, halo, (cyclo)alkyl, haloalkyl, (cyclo)alkoxy, CONR⁵R⁶, cycloalkylalkylene, cycloalkylalkoxy, or CO₂R⁷; R⁵ and R⁶ = independently H, alkyl, or taken together = unbranched alkylene; R⁷ = alkyl] were prep'd. as rotamase enzyme inhibitors, particularly FKBP-12 and FKBP-52 inhibitors. Thus, (2S)-1-(1,3-benzoxazol-2-yl)-2-piperidinecarboxylic acid (prepn. given) was amidated with (3S)-1-benzylpyrrolidine-3-ylamine in the presence of 1-hydroxybenzotriazole hydrate and 1-(3-dimethylaminopropyl)-3-ethylcarbodiimide.HCl in CH₂Cl₂ to yield II. Twenty-one compds. of the invention demonstrated inhibitory activity

against human recombinant FKBP-12 in a coupled colorimetric PPIase in vitro assay with IC50 values below 1200 nM, and II inhibited the rotamase enzyme FKBP-52 in a similar assay with IC50 = 2790 nM. As neurotrophic agents, the invention compds. promote neuronal regeneration and outgrowth and are useful for the treatment of neurodegenerative diseases or other disorders involving nerve damage.

RE.CNT 6 THERE ARE 6 CITED REFERENCES AVAILABLE FOR THIS RECORD
ALL CITATIONS AVAILABLE IN THE RE FORMAT

L6 ANSWER 4 OF 5 CAPLUS COPYRIGHT 2003 ACS on STN
AN 1998:719132 CAPLUS
DN 129:343719
TI Preparation of hemoregulatory peptides for stimulating the myelopoietic system
IN Bhatnagar, Pradip Kumar; Heerding, Dirk; Fischer, Peter Martin
PA Smithkline Beecham Corporation, USA; Nycomed Pharma As
SO U.S., 20 pp., Cont.-in-part of U.S. Ser. No. 66,952, abandoned.
CODEN: USXXAM
DT Patent
LA English
FAN.CNT 2

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
US 5830867	A	19981103	US 1996-553562	19960130
			US 1993-66952	B219930524
			US 1993-150524	B219931109
			WO 1994-US5859 W	19940524
WO 9427627	A1	19941208	WO 1994-US5859	19940524
W: AU, BB, BG, BR, BY, CA, CN, CZ, FI, HU, JP, KP, KR, KZ, LK, MG, MN, MW, NO, NZ, PL, RO, RU, SD, SI, SK, UA, US, US, VN				
RW: AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, ML, MR, NE, SN, TD, TG				
			US 1993-66952	A 19930524
			US 1993-150524	A 19931109
ZA 9403737	A	19950202	ZA 1994-3737	19940527
			US 1993-150524	19931109

PATENT FAMILY INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 9427627	A1	19941208	WO 1994-US5859	19940524
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			US 1993-66952	A 19930524
			US 1993-150524	A 19931109
AU 9470449	A1	19941220	AU 1994-70449	19940524
			US 1993-66952	A 19930524
			US 1993-150524	A 19931109
			WO 1994-US5859 W	19940524
EP 701446	A1	19960320	EP 1994-919232	19940524
R: BE, CH, DE, FR, GB, IT, LI, NL				
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			US 1993-150524	A 19931109
			WO 1994-US5859 W	19940524
JP 09504266	T2	19970428	JP 1995-500916	19940524
			US 1993-66952	A 19930524

			US 1993-150524 A 19931109
			WO 1994-US5859 W 19940524
ZA 9403737	A	19950202	ZA 1994-3737 19940527
			US 1993-150524 19931109
US 5830867	A	19981103	US 1996-553562 19960130
			US 1993-66952 B219930524
			US 1993-150524 B219931109
			WO 1994-US5859 W 19940524

OS MARPAT 129:343719

AB Peptides A1-B1-X1-(CH₂)_m-(CONR₁)_r-(CH₂)_s-Y1-(CH₂)_s-(NR₁CO)_r-(CH₂)_n-X1-B1-A1 [A1 = certain amino acids or aza heterocyclic acids; B1 = certain amino acids, 2-amino-3-hydroxythiopropionic acid, 2-amino-1-hydroxypropyl, or 2-amino-1-hydroxypent-3-enyl; X1 = O, S, NR₁, CR₂R₃; Y1 = O, S, NR₁, CR₂R₃, imidazolyl, triazolyl, Ph, etc.; R₁, R₂, R₃ = H, alkyl, imidazolyl, benzyl, etc.; m, n = 0-5; r = 0-2; s = 0, 1] were prepd. for stimulating the myelopoietic system. Thus, N,N'-bis(picolinoyl-seryl-.beta.-alanyl)-1,4-diaminobenzene (Boc = tert-butoxycarbonyl) was prepd. from 1,4-phenylenediamine dihydrochloride, Boc-.beta.-Ala-OH, Boc-Ser(Bzl)-OH, and picolinic acid.

RE.CNT 5 THERE ARE 5 CITED REFERENCES AVAILABLE FOR THIS RECORD
ALL CITATIONS AVAILABLE IN THE RE FORMAT

L6 ANSWER 5 OF 5 CAPLUS COPYRIGHT 2003 ACS on STN

AN 1993:552116 CAPLUS

DN 119:152116

TI Use of renin inhibitors for the treatment of glaucoma

IN Tanaka, Yoko; Kagayama, Akira; Hata, Takehisa

PA Fujisawa Pharmaceutical Co., Ltd., Japan

SO PCT Int. Appl., 25 pp.

CODEN: PIXXD2

DT Patent

LA English

FAN.CNT 1

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
PI	WO 9312796	A1	19930708	WO 1992-JP1656	19921218
	W: AU, CA, HU, JP, KR, RU, US				
	RW: AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE				
	ZA 9209738	A	19930617	GB 1991-27041	19911220
				ZA 1992-9738	19921215
				GB 1991-27041	19911220
	AU 9331712	A1	19930728	AU 1993-31712	19921218
	AU 661748	B2	19950803		
				GB 1991-27041	19911220
				WO 1992-JP1656	19921218
	EP 617622	A1	19941005	EP 1993-900396	19921218
	R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LI, LU, NL, PT, SE				
				GB 1991-27041	19911220
				WO 1992-JP1656	19921218
	JP 07506807	T2	19950727	JP 1992-511545	19921218
				GB 1991-27041	19911220
				WO 1992-JP1656	19921218
	CN 1088934	A	19940706	CN 1993-101190	19930102
				GB 1991-27041	19911220

OS MARPAT 119:152116

GI

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NEWS	6	AUG 18	Data available for download as a PDF in RDISCLOSURE
NEWS	7	AUG 18	Simultaneous left and right truncation added to PASCAL
NEWS	8	AUG 18	FROSTI and KOSMET enhanced with Simultaneous Left and Right Truncation
NEWS	9	AUG 18	Simultaneous left and right truncation added to ANABSTR
NEWS	10	SEP 22	DIPPR file reloaded
NEWS	11	SEP 25	INPADOC: Legal Status data to be reloaded
NEWS	12	SEP 29	DISSABS now available on STN
NEWS	13	OCT 10	PCTFULL: Two new display fields added
NEWS	14	OCT 21	BIOSIS file reloaded and enhanced
NEWS	15	OCT 28	BIOSIS file segment of TOXCENTER reloaded and enhanced
NEWS EXPRESS		OCTOBER 01	CURRENT WINDOWS VERSION IS V6.01a, CURRENT MACINTOSH VERSION IS V6.0b(ENG) AND V6.0Jb(JP), AND CURRENT DISCOVER FILE IS DATED 23 SEPTEMBER 2003
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Patel

<11/4/2003>

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DICTIONARY FILE UPDATES: 3 NOV 2003 HIGHEST RN 612478-18-9

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conducting SmartSELECT searches.

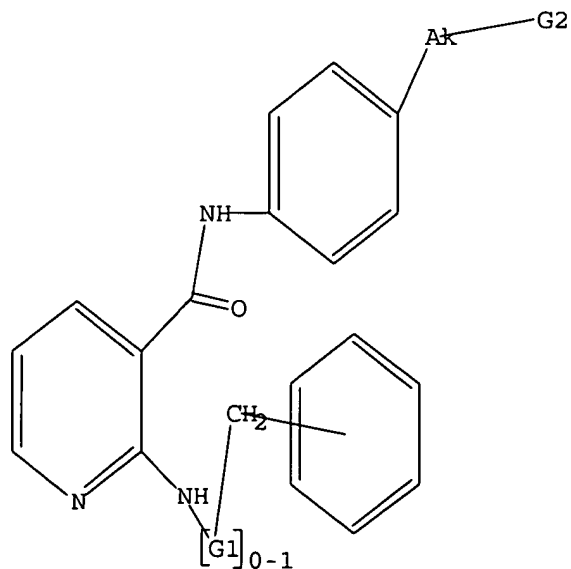
Crossover limits have been increased. See HELP CROSSOVER for details.

Experimental and calculated property data are now available. See HELP
PROPERTIES for more information. See STNote 27, Searching Properties
in the CAS Registry File, for complete details:
<http://www.cas.org/ONLINE/STN/STNOTES/stnotes27.pdf>

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L1 STRUCTURE UPLOADED

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L1 HAS NO ANSWERS
L1 STR



G1 S, S02

G2 Cb, Cy, Hy

Structure attributes must be viewed using STN Express query preparation.

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FULL SEARCH INITIATED 14:58:46 FILE 'REGISTRY'

FULL SCREEN SEARCH COMPLETED - 779 TO ITERATE

100.0% PROCESSED 779 ITERATIONS

2 ANSWERS

SEARCH TIME: 00.00.01

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=> file caplus

COST IN U.S. DOLLARS

SINCE FILE

TOTAL

ENTRY

SESSION

FULL ESTIMATED COST

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FILE COVERS 1907 - 4 Nov 2003 VOL 139 ISS 19

FILE LAST UPDATED: 3 Nov 2003 (20031103/ED)

This file contains CAS Registry Numbers for easy and accurate substance identification.

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L3 2 L2

=> d l3 fbib hitstr abs total

L3 ANSWER 1 OF 2 CAPLUS COPYRIGHT 2003 ACS on STN

AN 2003:551181 CAPLUS

DN 139:117339

TI Preparation of substituted arylamine derivatives as antitumor agents

IN Elbaum, Daniel; Askew, Benny; Booker, Shon; Germain, Julie; Habgood, Gregory; Handley, Michael; Kim, Tae-Seong; Li, Aiwen; Nishimura, Nobuko; Patel, Vinod F.; Yuan, Chester Chenguang; Kim, Joseph L.

PA Amgen Inc., USA

SO U.S. Pat. Appl. Publ., 106 pp., Cont.-in-part of U.S. Ser. No. 46,526.

CODEN: USXXCO

DT Patent

LA English

FAN.CNT 2

PATENT NO.

KIND DATE

APPLICATION NO. DATE

Patel

<11/4/2003>

PI	US 2003134836	A1	20030717	US 2002-197960	20020717
				US 2001-261360PP	20010112
				US 2001-323686PP	20010919
	US 2002147198	A1	20021010	US 2002-46526 A2	20020110
				US 2002-46526	20020110
				US 2001-261360PP	20010112
				US 2001-323686PP	20010919

PATENT FAMILY INFORMATION:

FAN 2002:539663

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
PI	WO 2002055501	A2	20020718	WO 2002-US742	20020111
	WO 2002055501	A3	20021219		
	W:	AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, OM, PH, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TN, TR, TT, TZ, UA, UG, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM			
	RW:	GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG			
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				US 2001-323686PP	20010919
				US 2002-46526 A	20020110
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				US 2001-261360PP	20010112
				US 2001-323686PP	20010919

OS MARPAT 139:117339

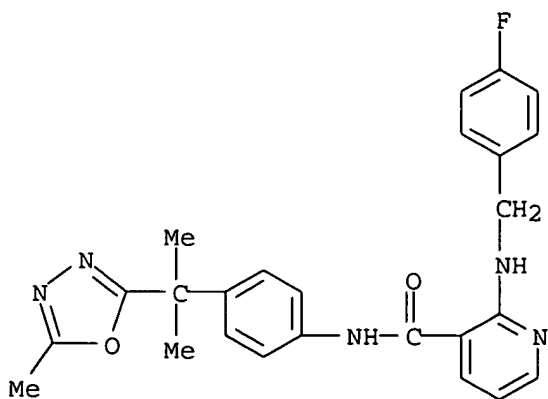
IT **561297-64-1P**

RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(prepn. of substituted aminopyridines as antitumor agents)

RN 561297-64-1 CAPLUS

CN 3-Pyridinecarboxamide, 2-[[[4-(4-fluorophenyl)methyl]amino]-N-[4-[1-methyl-1-(5-methyl-1,3,4-oxadiazol-2-yl)ethyl]phenyl]- (9CI) (CA INDEX NAME)

IT **442846-35-7P**

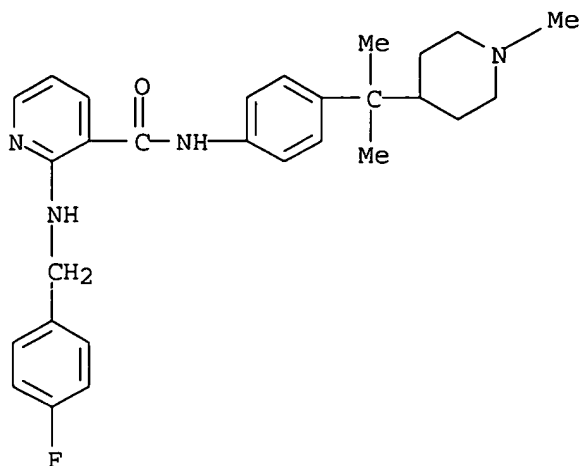
RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU

(Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

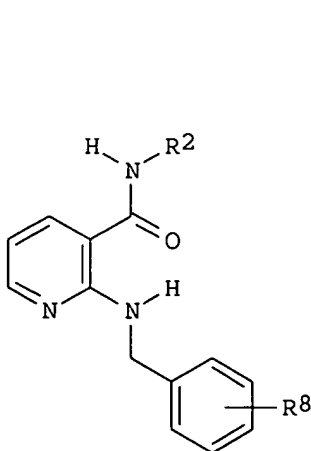
(target compd.; prepn. of substituted aminopyridines as antitumor agents)

RN 442846-35-7 CAPLUS

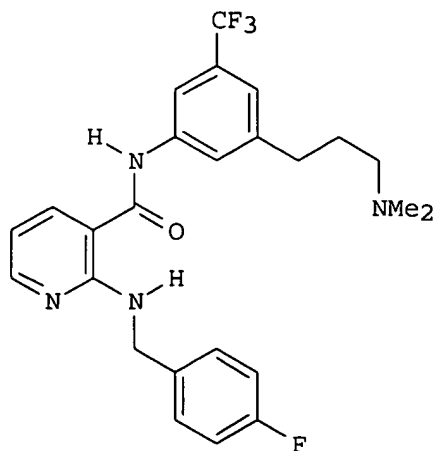
CN 3-Pyridinecarboxamide, 2-[[[4-(4-fluorophenyl)methyl]amino]-N-[4-[1-methyl-1-(1-methyl-4-piperidinyl)ethyl]phenyl]- (9CI) (CA INDEX NAME)



GI



I



II

AB The title compds. I [R2 = (un)substituted Ph, 9-10 membered bicyclic and 11-14 membered tricyclic (un)satd. heterocyclyl; R8 = halo, NH2, NO2, etc.], and their pharmaceutically acceptable derivs., are prepd. and disclosed as agents effective for prophylaxis and treatment of diseases, such as angiogenesis mediated diseases. E.g., a multi-step synthesis of II, starting from 1-dimethylamino-2-propyne and 3-bromo-5-trifluoromethylaniline, was given. Selected compds. of the invention,

e.g., II, inhibited VEGF-stimulated cell proliferation at a level below 50 nM. The invention encompasses novel compds., analogs, prodrugs and pharmaceutically acceptable derivs. thereof, pharmaceutical compns. and methods for prophylaxis and treatment of diseases and other maladies or conditions involving, cancer and the like.

L3 ANSWER 2 OF 2 CAPLUS COPYRIGHT 2003 ACS on STN

AN 2002:539663 CAPLUS

DN 137:109210

TI Preparation of substituted arylamine derivatives and methods of use as antitumor agents

IN Chen, Guoqing; Booker, Shon; Cai, Guolin; Croghan, Michael; Dipietro, Lucian; Dominguez, Celia; Elbaum, Daniel; Germain, Julie; Huang, Qi; Kim, Joseph L.; Kim, Tae-Seong; Patel, Vinod F.; Smith, Leon M.; Tasker, Andrew; Xi, Ning; Xu, Shimin; Yuan, Chester Chenguang

PA Amgen Inc., USA

SO PCT Int. Appl., 253 pp.

CODEN: PIXXD2

DT Patent

LA English

FAN.CNT 2

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
PI	WO 2002055501	A2	20020718	WO 2002-US742	20020111
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	RW:				
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				US 2001-323686PP	20010919
				US 2002-46526 A	20020110
	US 2002147198	A1	20021010	US 2002-46526	20020110
				US 2001-261360PP	20010112
				US 2001-323686PP	20010919

PATENT FAMILY INFORMATION:

FAN 2003:551181

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
PI	US 2003134836	A1	20030717	US 2002-197960	20020717
				US 2001-261360PP	20010112
				US 2001-323686PP	20010919
				US 2002-46526 A2	20020110
	US 2002147198	A1	20021010	US 2002-46526	20020110
				US 2001-261360PP	20010112
				US 2001-323686PP	20010919

OS MARPAT 137:109210

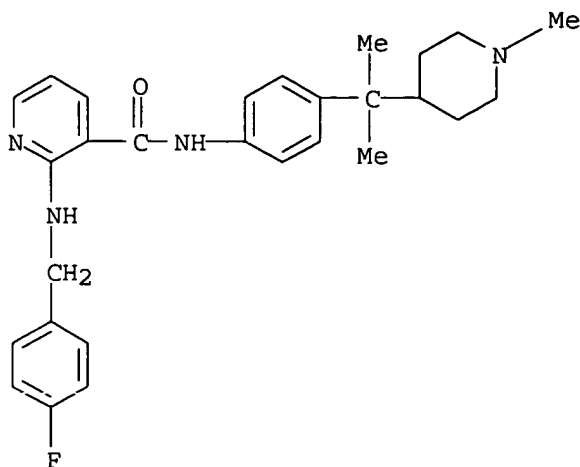
IT 442846-35-7P

RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(target compd.; prepn. of substituted aminopyridines as antitumor agents)

RN 442846-35-7 CAPLUS

CN 3-Pyridinecarboxamide, 2-[[[(4-fluorophenyl)methyl]amino]-N-[4-[1-methyl-1-(1-methyl-4-piperidiny)ethyl]phenyl]]- (9CI) (CA INDEX NAME)



GI

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

AB Title compds. I [B1 and B2 independently equal C or N, wherein B1B2 form part of 5-6 membered heteroaryl ring A; R1 = one or more substituents selected from H, halo, oxo, (un)substituted cycloalkyl, phenylalkyl, etc.; R2 = (un)substituted cycloalkyl, cycloalkenyl, 6-10 membered aryl or 5-6 membered heterocyclyl, etc.; R3 = (un)substituted aryl; R4 = H, alkyl, (un)substituted Ph or aralkyl; X1 = bond, alkylene, alkenylene and alkynylene, where one of the CH2 groups may be substituted with O or NH, wherein X1 is optionally substituted with OH; X2 = (un)substituted N contg. linker, e.g., -NHCH2-], and there pharmaceutically acceptable derivs., are prep'd. and disclosed as agents effective for prophylaxis and treatment of diseases, such as angiogenesis mediated diseases. Thus, II was prep'd. via arylation of 1-dimethylamino-2-propyne with 3-bromo-5-trifluoromethylaniline, hydrogenation, amidation with 2-chloropyridine-3-carbonyl chloride and chloro-substitution with 4-fluorobenzylamine. Selected compds. of the invention, e.g., II, inhibited VEGF-stimulated cell proliferation at a level below 50 nM. The invention encompasses novel compds., analogs, prodrugs and pharmaceutically acceptable derivs. thereof, pharmaceutical compns. and methods for prophylaxis and treatment of diseases and other maladies or conditions involving, cancer and the like.

=> log y

COST IN U.S. DOLLARS

SINCE FILE

TOTAL

ENTRY

SESSION

FULL ESTIMATED COST

21.56

169.92

DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS)

SINCE FILE

TOTAL

Patel

<11/4/2003>

	ENTRY	SESSION
CA SUBSCRIBER PRICE	-1.30	-1.30

STN INTERNATIONAL LOGOFF AT 14:59:18 ON 04 NOV 2003

Welcome to STN International! Enter x:x

LOGINID:sssptal611sxp

PASSWORD:

* * * * * RECONNECTED TO STN INTERNATIONAL * * * * *

SESSION RESUMED IN FILE 'HOME' AT 13:18:23 ON 30 JUN 2003

FILE 'HOME' ENTERED AT 13:18:23 ON 30 JUN 2003

COST IN U.S. DOLLARS

SINCE FILE

TOTAL

ENTRY

SESSION

FULL ESTIMATED COST

0.63

0.63

=> file reg

COST IN U.S. DOLLARS

SINCE FILE

TOTAL

ENTRY

SESSION

FULL ESTIMATED COST

0.63

0.63

FILE 'REGISTRY' ENTERED AT 13:18:37 ON 30 JUN 2003

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STRUCTURE FILE UPDATES: 29 JUN 2003 HIGHEST RN 539790-82-4

DICTIONARY FILE UPDATES: 29 JUN 2003 HIGHEST RN 539790-82-4

TSCA INFORMATION NOW CURRENT THROUGH JANUARY 6, 2003

Please note that search-term pricing does apply when conducting SmartSELECT searches.

Crossover limits have been increased. See HELP CROSSOVER for details.

Experimental and calculated property data are now available. See HELP PROPERTIES for more information. See STNote 27, Searching Properties in the CAS Registry File, for complete details:
<http://www.cas.org/ONLINE/STN/STNOTES/stnotes27.pdf>

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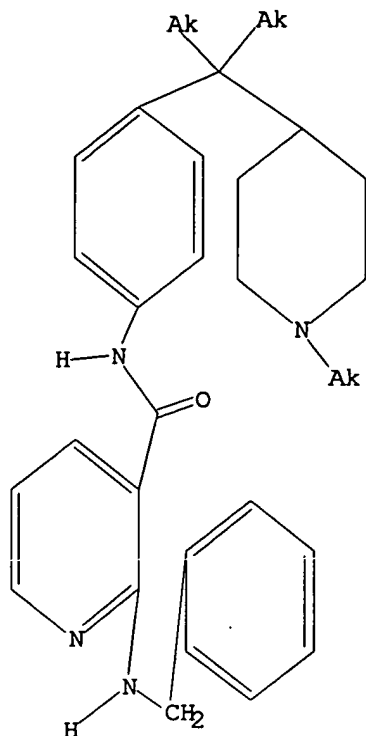
Uploading 10046526.1

L1 STRUCTURE UPLOADED

=> d l1

L1 HAS NO ANSWERS

L1 STR



Structure attributes must be viewed using STN Express query preparation.

=> s l1

SAMPLE SEARCH INITIATED 13:19:04 FILE 'REGISTRY'
 SAMPLE SCREEN SEARCH COMPLETED - 7 TO ITERATE

100.0% PROCESSED 7 ITERATIONS 0 ANSWERS
 SEARCH TIME: 00.00.01

FULL FILE PROJECTIONS: ONLINE **COMPLETE**
 BATCH **COMPLETE**
 PROJECTED ITERATIONS: 7 TO 298
 PROJECTED ANSWERS: 0 TO 0

L2 0 SEA SSS SAM L1

=> s l1 sss full

FULL SEARCH INITIATED 13:19:12 FILE 'REGISTRY'
 FULL SCREEN SEARCH COMPLETED - 85 TO ITERATE

100.0% PROCESSED 85 ITERATIONS 1 ANSWERS
 SEARCH TIME: 00.00.01

L3 1 SEA SSS FUL L1

=> file caplus

COST IN U.S. DOLLARS

SINCE FILE

TOTAL

Patel

<11/4/2003>

	ENTRY	SESSION
FULL ESTIMATED COST	148.15	148.78

FILE 'CAPLUS' ENTERED AT 13:19:17 ON 30 JUN 2003
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FILE COVERS 1907 - 30 Jun 2003 VOL 139 ISS 1
 FILE LAST UPDATED: 29 Jun 2003 (20030629/ED)

This file contains CAS Registry Numbers for easy and accurate substance identification.

=> s 13

L4 1 L3

=> d 14 fbib hitstr abs total

L4 ANSWER 1 OF 1 CAPLUS COPYRIGHT 2003 ACS
 AN 2002:539663 CAPLUS
 DN 137:109210
 TI Preparation of substituted arylamine derivatives and methods of use as antitumor agents
 IN Chen, Guoqing; Booker, Shon; Cai, Guolin; Croghan, Michael; Dipietro, Lucian; Dominguez, Celia; Elbaum, Daniel; Germain, Julie; Huang, Qi; Kim, Joseph L.; Kim, Tae-Seong; Patel, Vinod F.; Smith, Leon M.; Tasker, Andrew; Xi, Ning; Xu, Shimin; Yuan, Chester Chenguang
 PA Amgen Inc., USA
 SO PCT Int. Appl., 253 pp.
 CODEN: PIXXD2
 DT Patent
 LA English
 FAN.CNT 1

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
PI	WO 2002055501	A2	20020718	WO 2002-US742	20020111
	WO 2002055501	A3	20021219		
W:	AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, OM, PH, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TN, TR, TT, TZ, UA, UG, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM				
RW:	GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR,				

Patel

<11/4/2003>

BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG

US 2001-261360PP 20010112

US 2001-323686PP 20010919

US 2002-46526 A 20020110

US 2002147198 A1 20021010

US 2002-46526 20020110

US 2001-261360PP 20010112

US 2001-323686PP 20010919

OS MARPAT 137:109210

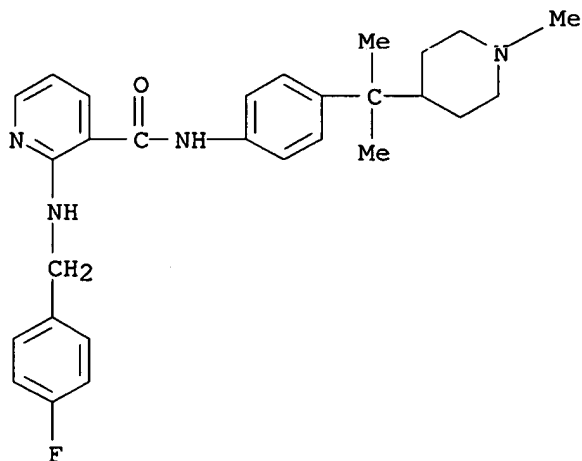
IT 442846-35-7P

RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU
(Therapeutic use); BIOL (Biological study); PREP (Preparation); USES
(Uses)

(target compd.; prepn. of substituted aminopyridines as antitumor
agents)

RN 442846-35-7 CAPLUS

CN 3-Pyridinecarboxamide, 2-[[[(4-fluorophenyl)methyl]amino]-N-[4-[1-methyl-1-(1-methyl-4-piperidinyl)ethyl]phenyl]- (9CI) (CA INDEX NAME)



GI

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

AB Title compds. I [B1 and B2 independently equal C or N, wherein B1B2 form part of 5-6 membered heteroaryl ring A; R1 = one or more substituents selected from H, halo, oxo, (un)substituted cycloalkyl, phenylalkyl, etc.; R2 = (un)substituted cycloalkyl, cycloalkenyl, 6-10 membered aryl or 5-6 membered heterocyclyl, etc.; R3 = (un)substituted aryl; R4 = H, alkyl, (un)substituted Ph or aralkyl; X1 = bond, alkylene, alkenylene and alkynylene, where one of the CH2 groups may be substituted with O or NH, wherein X1 is optionally substituted with OH; X2 = (un)substituted N contg. linker, e.g., -NHCH2-], and there pharmaceutically acceptable derivs., are prepd. and disclosed as agents effective for prophylaxis and treatment of diseases, such as angiogenesis mediated diseases. Thus, II was prepd. via arylation of 1-dimethylamino-2-propyne with 3-bromo-5-trifluoromethylaniline, hydrogenation, amidation with 2-chloropyridine-3-carbonyl chloride and chloro-substitution with

4-phenoxyaniline. Selected compds. of the invention, e.g., II, inhibited VEGF-stimulated cell proliferation at a level below 50 nM. The invention encompasses novel compds., analogs, prodrugs and pharmaceutically acceptable derivs. thereof, pharmaceutical compns. and methods for prophylaxis and treatment of diseases and other maladies or conditions involving, cancer and the like.

=> d cost

COST IN U.S. DOLLARS

	SINCE FILE ENTRY	TOTAL SESSION
CONNECT CHARGES	0.34	1.13
NETWORK CHARGES	0.06	0.30
SEARCH CHARGES	0.00	147.75
DISPLAY CHARGES	4.32	4.32
	-----	-----
	4.72	153.50
CAPLUS FEE (5%)	0.23	0.23
	-----	-----
FULL ESTIMATED COST	4.95	153.73

DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS)

	SINCE FILE ENTRY	TOTAL SESSION
CA SUBSCRIBER PRICE	-0.65	-0.65

IN FILE 'CAPLUS' AT 13:19:53 ON 30 JUN 2003

=> log y

COST IN U.S. DOLLARS

	SINCE FILE ENTRY	TOTAL SESSION
FULL ESTIMATED COST	5.79	154.57
DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS)		
	SINCE FILE ENTRY	TOTAL SESSION
CA SUBSCRIBER PRICE	-0.65	-0.65

STN INTERNATIONAL LOGOFF AT 13:21:02 ON 30 JUN 2003

Welcome to STN International! Enter x:x

LOGINID:sssptal611sxp

PASSWORD:

TERMINAL (ENTER 1, 2, 3, OR ?):2

* * * * * Welcome to STN International * * * * *

NEWS	1		Web Page URLs for STN Seminar Schedule - N. America
NEWS	2		"Ask CAS" for self-help around the clock
NEWS	3	Jun 03	New e-mail delivery for search results now available
NEWS	4	Aug 08	PHARMAMarketLetter(PHARMAML) - new on STN
NEWS	5	Aug 19	Aquatic Toxicity Information Retrieval (AQUIRE) now available on STN
NEWS	6	Aug 26	Sequence searching in REGISTRY enhanced
NEWS	7	Sep 03	JAPIO has been reloaded and enhanced
NEWS	8	Sep 16	Experimental properties added to the REGISTRY file
NEWS	9	Sep 16	CA Section Thesaurus available in CAPLUS and CA
NEWS	10	Oct 01	CASREACT Enriched with Reactions from 1907 to 1985
NEWS	11	Oct 24	BEILSTEIN adds new search fields
NEWS	12	Oct 24	Nutraceuticals International (NUTRACEUT) now available on STN
NEWS	13	Nov 18	DKILIT has been renamed APOLLIT
NEWS	14	Nov 25	More calculated properties added to REGISTRY
NEWS	15	Dec 04	CSA files on STN
NEWS	16	Dec 17	PCTFULL now covers WP/PCT Applications from 1978 to date
NEWS	17	Dec 17	TOXCENTER enhanced with additional content
NEWS	18	Dec 17	Adis Clinical Trials Insight now available on STN
NEWS	19	Jan 29	Simultaneous left and right truncation added to COMPENDEX, ENERGY, INSPEC
NEWS	20	Feb 13	CANCERLIT is no longer being updated
NEWS	21	Feb 24	METADEx enhancements
NEWS	22	Feb 24	PCTGEN now available on STN
NEWS	23	Feb 24	TEMA now available on STN
NEWS	24	Feb 26	NTIS now allows simultaneous left and right truncation
NEWS	25	Feb 26	PCTFULL now contains images
NEWS	26	Mar 04	SDI PACKAGE for monthly delivery of multifile SDI results
NEWS	27	Mar 20	EVENTLINE will be removed from STN
NEWS	28	Mar 24	PATDPAFULL now available on STN
NEWS	29	Mar 24	Additional information for trade-named substances without structures available in REGISTRY
NEWS	30	Apr 11	Display formats in DGENE enhanced
NEWS	31	Apr 14	MEDLINE Reload
NEWS	32	Apr 17	Polymer searching in REGISTRY enhanced
NEWS	33	Jun 13	Indexing from 1947 to 1956 added to records in CA/CAPLUS
NEWS	34	Apr 21	New current-awareness alert (SDI) frequency in WPIDS/WPINDEX/WPIX
NEWS	35	Apr 28	RDISCLOSURE now available on STN
NEWS	36	May 05	Pharmacokinetic information and systematic chemical names added to PHAR
NEWS	37	May 15	MEDLINE file segment of TOXCENTER reloaded
NEWS	38	May 15	Supporter information for ENCOMPPAT and ENCOMPLIT updated
NEWS	39	May 16	CHEMREACT will be removed from STN
NEWS	40	May 19	Simultaneous left and right truncation added to WSCA

NEWS 41 May 19 RAPRA enhanced with new search field, simultaneous left and right truncation
NEWS 42 Jun 06 Simultaneous left and right truncation added to CBNB
NEWS 43 Jun 06 PASCAL enhanced with additional data
NEWS 44 Jun 20 2003 edition of the FSTA Thesaurus is now available
NEWS 45 Jun 25 HSDB has been reloaded

NEWS EXPRESS April 4 CURRENT WINDOWS VERSION IS V6.01a, CURRENT
MACINTOSH VERSION IS V6.0b(ENG) AND V6.0Jb(JP),
AND CURRENT DISCOVER FILE IS DATED 01 APRIL 2003
NEWS HOURS STN Operating Hours Plus Help Desk Availability
NEWS INTER General Internet Information
NEWS LOGIN Welcome Banner and News Items
NEWS PHONE Direct Dial and Telecommunication Network Access to STN
NEWS WWW CAS World Wide Web Site (general information)

Enter NEWS followed by the item number or name to see news on that specific topic.

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* * * * * STN Columbus * * * * *

FILE 'HOME' ENTERED AT 13:25:55 ON 30 JUN 2003

=> file reg

COST IN U.S. DOLLARS

SINCE FILE	TOTAL
ENTRY	SESSION
0.21	0.21

FULL ESTIMATED COST

FILE 'REGISTRY' ENTERED AT 13:26:04 ON 30 JUN 2003

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STRUCTURE FILE UPDATES: 29 JUN 2003 HIGHEST RN 539790-82-4

DICTIONARY FILE UPDATES: 29 JUN 2003 HIGHEST RN 539790-82-4

TSCA INFORMATION NOW CURRENT THROUGH JANUARY 6, 2003

Please note that search-term pricing does apply when conducting SmartSELECT searches.

Crossover limits have been increased. See HELP CROSSOVER for details.

Experimental and calculated property data are now available. See HELP PROPERTIES for more information. See STNote 27, Searching Properties in the CAS Registry File, for complete details:
<http://www.cas.org/ONLINE/STN/STNOTES/stnotes27.pdf>

=>

Patel

<11/4/2003>

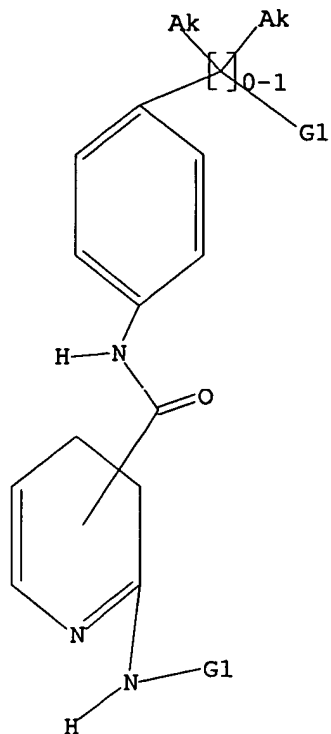
Uploading 10046526.2

L1 STRUCTURE UPLOADED

=> d l1

L1 HAS NO ANSWERS

L1 STR



G1 Cb,Cy,Hy,Ak

Structure attributes must be viewed using STN Express query preparation.

=> s l1

SAMPLE SEARCH INITIATED 13:26:26 FILE 'REGISTRY'

SAMPLE SCREEN SEARCH COMPLETED - 2066 TO ITERATE

48.4% PROCESSED 1000 ITERATIONS
INCOMPLETE SEARCH (SYSTEM LIMIT EXCEEDED)
SEARCH TIME: 00.00.01

10 ANSWERS

FULL FILE PROJECTIONS: ONLINE **COMPLETE**
BATCH **COMPLETE**
PROJECTED ITERATIONS: 38594 TO 44046
PROJECTED ANSWERS: 141 TO 685

L2 10 SEA SSS SAM L1

=> s l1 sss full

FULL SEARCH INITIATED 13:26:33 FILE 'REGISTRY'

FULL SCREEN SEARCH COMPLETED - 40731 TO ITERATE

Patel

<11/4/2003>

100.0% PROCESSED 40731 ITERATIONS
SEARCH TIME: 00.00.02

255 ANSWERS

L3 255 SEA SSS FUL L1

=> file caplus

COST IN U.S. DOLLARS

SINCE FILE

TOTAL

ENTRY

SESSION

FULL ESTIMATED COST

148.15

148.36

FILE 'CAPLUS' ENTERED AT 13:26:41 ON 30 JUN 2003

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FILE COVERS 1907 - 30 Jun 2003 VOL 139 ISS 1

FILE LAST UPDATED: 29 Jun 2003 (20030629/ED)

This file contains CAS Registry Numbers for easy and accurate substance identification.

=> s l3

L4 20 L3

=> d l4 fbib hitstr abs total

L4 ANSWER 1 OF 20 CAPLUS COPYRIGHT 2003 ACS

AN 2003:319701 CAPLUS

DN 138:337840

TI Preparation of 5'-acylamino-1,1'-biphenyl-4-carboxamides as p38 kinase inhibitors

IN Angell, Richard Martyn; Aston, Nicola Mary; Bamborough, Paul; Bamford, Mark James; Cockerill, George Stuart; Flack, Stephen Sean; Laine, Dramane Ibrahim; Merrick, Suzanne Joy; Smith, Kathryn Jane; Walker, Ann Louise

PA Glaxo Group Limited, UK

SO PCT Int. Appl., 64 pp.

CODEN: PIXXD2

DT Patent

LA English

FAN.CNT 1

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
PI	WO 2003032971	A1	20030424	WO 2002-EP11576	20021016
	W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH,				

Patel

<11/4/2003>

GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR,
 LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, OM, PH,
 PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TN, TR, TT, TZ,
 UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW, AM, AZ, BY, KG, KZ, MD,
 RU, TJ, TM

RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AT, BE, BG,
 CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL,
 PT, SE, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR,
 NE, SN, TD, TG

GB 2001-24939 A 20011017

OS MARPAT 138:337840

IT 515812-31-4P 515812-32-5P 515812-34-7P

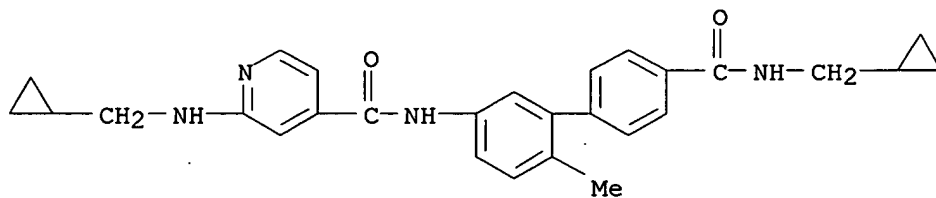
515812-35-8P 515812-44-9P

RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU
 (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES
 (Uses)

(prepn. of 5'-acylamino-1,1'-biphenyl-4-carboxamides as p38 kinase
 inhibitors)

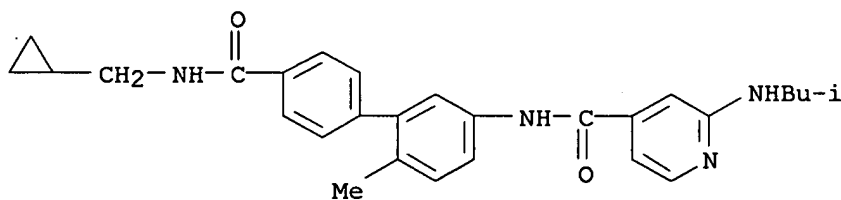
RN 515812-31-4 CAPLUS

CN 4-Pyridinecarboxamide, 2-[(cyclopropylmethyl)amino]-N-[4'-
 [[(cyclopropylmethyl)amino]carbonyl]-6-methyl[1,1'-biphenyl]-3-yl]- (9CI)
 (CA INDEX NAME)



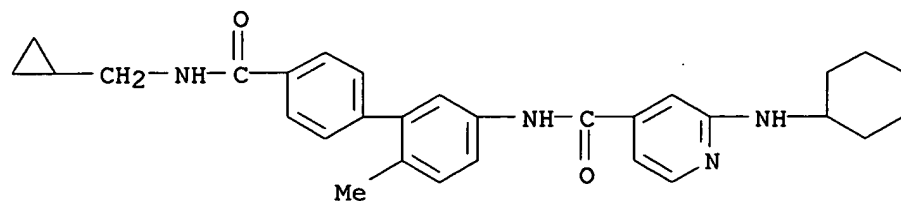
RN 515812-32-5 CAPLUS

CN 4-Pyridinecarboxamide, N-[4'-[[[(cyclopropylmethyl)amino]carbonyl]-6-
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 NAME)



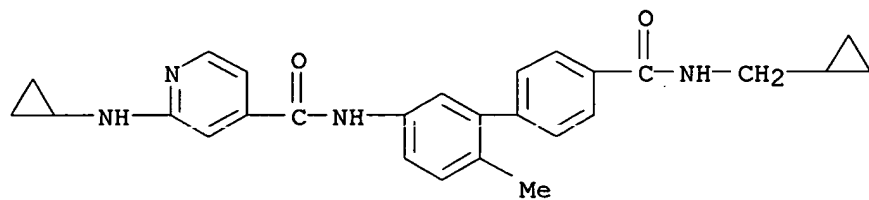
RN 515812-34-7 CAPLUS

CN 4-Pyridinecarboxamide, 2-(cyclohexylamino)-N-[4'-
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 (CA INDEX NAME)



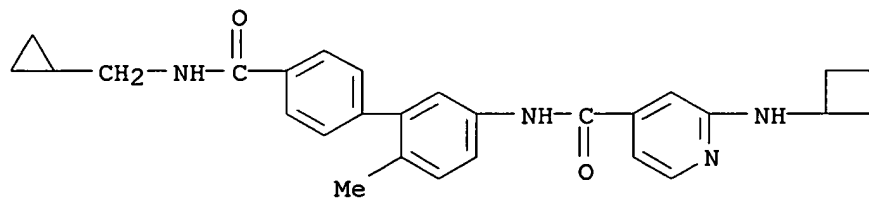
RN 515812-35-8 CAPLUS

CN 4-Pyridinecarboxamide, 2-(cyclopropylamino)-N-[4'-
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 (CA INDEX NAME)

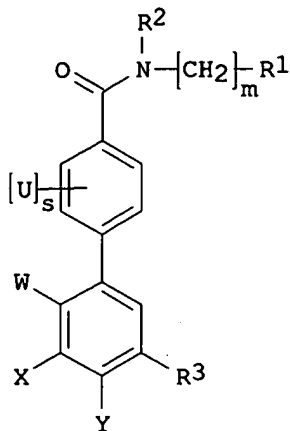


RN 515812-44-9 CAPLUS

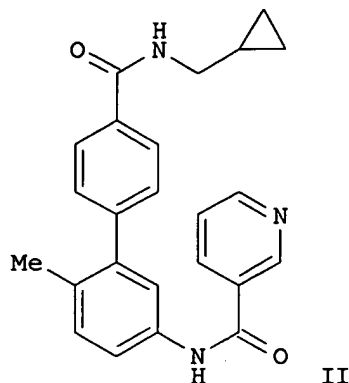
CN 4-Pyridinecarboxamide, 2-(cyclobutylamino)-N-[4'-
 [(cyclobutylmethyl)amino]carbonyl]-6-methyl[1,1'-biphenyl]-3-yl]- (9CI)
 (CA INDEX NAME)



GI



I



II

AB The title compds. [I; when m = 0-4, R1 = alkyl, cycloalkyl, alkenyl, etc.; when m = 2-4, R1 addnl. = alkoxy, OH, etc.; R2 = H, alkyl, (CH2)ncycloalkyl; R3 = NHCOR6 (wherein R6 = H, alkyl, alkoxy, etc.); U = Me, halo; W = Me, Cl; X, Y = H, Me, halo; m = 0-4; n = 0-1; s = 0-2], useful as pharmaceuticals, particularly as p38 kinase inhibitors, were prepd. E.g., a 6-step synthesis of the nicotinamide II, starting with 3-bromo-4-methylaniline, was given.

RE.CNT 2 THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS RECORD
ALL CITATIONS AVAILABLE IN THE RE FORMAT

L4 ANSWER 2 OF 20 CAPLUS COPYRIGHT 2003 ACS

AN 2003:261670 CAPLUS

DN 138:287666

TI Preparation of heteroaryllactams as Factor Xa inhibitors

IN Pinto, Donald; Quan, Mimi; Orwat, Michael; Li, Yun-Long; Han, Wei; Qiao, Jennifer; Lam, Patrick; Koch, Stephanie

PA Bristol-Myers Squibb Company, USA

SO PCT Int. Appl., 441 pp.

CODEN: PIXXD2

DT Patent

LA English

FAN.CNT 1

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 2003026652	A1	20030403	WO 2002-US29491	20020917
<p>W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, OM, PH, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM</p> <p>RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG</p>				

US 2001-324165PP 20010921

OS MARPAT 138:287666

IT 503613-25-OP, 2-[(4-Chlorobenzoyl)amino]-N-[4-(2-oxo-1-

Patel

<11/4/2003>

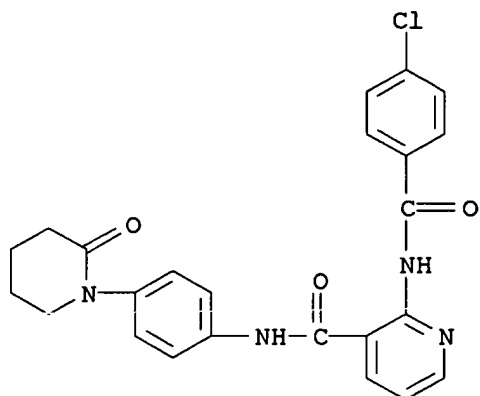
piperidinyl)phenyl]nicotinamide

RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(claimed compd.; prepn. of heteroaryllactams as Factor Xa inhibitors)

RN 503613-25-0 CAPLUS

CN 3-Pyridinecarboxamide, 2-[(4-chlorobenzoyl)amino]-N-[4-(2-oxo-1-piperidinyl)phenyl]- (9CI) (CA INDEX NAME)



AB P4PMM4 [M = 3-10 membered (substituted) (unsatd.) carbocyclyl, 4-10 membered heterocyclyl; P = null, 5-7 membered (substituted) (unsatd.) carbocyclyl, heterocyclyl fused to ring M; 1 of P4, M4 = ZAB, the other = G1G; G = (benzo-, pyrido-, pyrimido-, pyrazino-, or pyridazino-fused) (substituted) (unsatd.) 5-6 membered (hetero)cyclyl; G1 = null, (CR3R3a)1-5, etc.; R3, R3a = H, Me, Et, Pr, Ph, PhCH2, etc.; Z = bond, (CR3R3e)1-4, etc.; R3e = H, SO2NHR3, SO2N(R3)2, COR3, (substituted) alkyl, alkenyl, alkynyl, etc.; A = (substituted) 3-10 membered carbocyclyl, 5-12 membered heterocyclyl; Z = XNQ; X = null, CO, SO, SO2, etc.; NQ = 4-8 membered mono- or bicyclic (substituted) (unsatd.) ring contg. a CO or SO2 group adjacent to the N atom; with provisos], were prepd. Thus, 6-(4-iodophenyl)-3-methoxy-1-(4-methoxyphenyl)-1,4,5,6-tetrahydro-7H-pyrazolo[3,4-c]pyridin-7-one (prepn. given), .delta.-valerolactam, K2CO3, and CuI were refluxed in Me2SO to give 15% 3-methoxy-1-(4-methoxyphenyl)-6-[4-(2-oxo-1-piperidinyl)phenyl]-1,4,5,6-tetrahydro-7H-pyrazolo[3,4-c]pyridin-7-one. Several title compds. inhibited Factor Xa with IC50.ltoreq. 10 .mu.M.

RE.CNT 1 THERE ARE 1 CITED REFERENCES AVAILABLE FOR THIS RECORD
ALL CITATIONS AVAILABLE IN THE RE FORMAT

L4 ANSWER 3 OF 20 CAPLUS COPYRIGHT 2003 ACS

AN 2003:22869 CAPLUS

DN 138:89806

TI Preparation of arylpyrazoles as soluble epoxide hydrolase inhibitors for treatment of cardiovascular disease.

IN Ingraham, Richard H.; Proudfoot, John R.

PA Boehringer Ingelheim Pharmaceuticals Inc., USA

SO PCT Int. Appl., 44 pp.

CODEN: PIXXD2

DT Patent

LA English

FAN.CNT 1